

# Machine Learning Accelerates the Discovery of Design Rules and Exceptions in Stable Metal–Oxo Intermediate Formation

Aditya Nandy,<sup>†,‡</sup> Jiazhou Zhu,<sup>§</sup> Jon Paul Janet,<sup>†</sup> Chenru Duan,<sup>†,‡</sup> Rachel B. Getman,<sup>\*,§</sup> and Heather J. Kulik<sup>\*,†</sup>

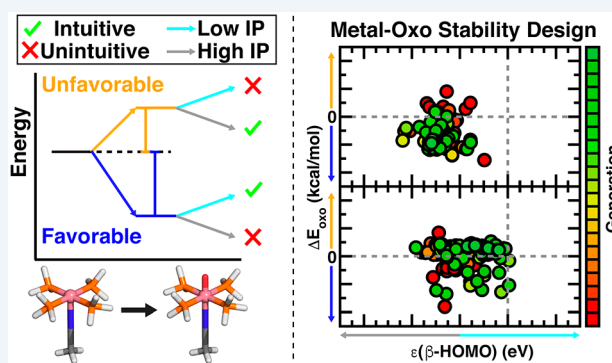
<sup>†</sup>Department of Chemical Engineering and <sup>‡</sup>Department of Chemistry, Massachusetts Institute of Technology, Cambridge, Massachusetts 02139, United States

<sup>§</sup>Department of Chemical & Biomolecular Engineering, Clemson University, Clemson, South Carolina 29634, United States

## S Supporting Information

**ABSTRACT:** Metal–oxo moieties are important catalytic intermediates in the selective partial oxidation of hydrocarbons and in water splitting. Stable metal–oxo species have reactive properties that vary depending on the spin state of the metal, complicating the development of structure–property relationships. To overcome these challenges, we train machine-learning (ML) models capable of predicting metal–oxo formation energies across a range of first-row metals, oxidation states, and spin states. Using connectivity-only features tailored for inorganic chemistry as inputs to kernel ridge regression or artificial neural network (ANN) ML models, we achieve good mean absolute errors (4–5 kcal/mol) on set-aside test data across a range of ligand orientations. Analysis of feature importance for oxo formation energy prediction reveals the dominance of nonlocal, electronic ligand properties in contrast to other transition metal complex properties (e.g., spin-state or ionization potential). We enumerate the theoretical catalyst space with an ANN, revealing expected trends in oxo formation energetics, such as destabilization of the metal–oxo species with increasing d-filling, as well as exceptions, such as weak correlations with indicators of oxidative stability of the metal in the resting state or unexpected spin-state dependence in reactivity. We carry out uncertainty-aware evolutionary optimization using the ANN to explore a >37 000 candidate catalyst space. New metal and oxidation state combinations are uncovered and validated with density functional theory (DFT), including counterintuitive oxo formation energies for oxidatively stable complexes. This approach doubles the density of confirmed DFT leads in originally sparsely populated regions of property space, highlighting the potential of ML-model-driven discovery to uncover catalyst design rules and exceptions.

**KEYWORDS:** metal–oxo species, machine learning, density functional theory, spin-state-dependent reactivity, transition metal catalysis, artificial neural networks



## 1. INTRODUCTION

The selective partial oxidation of alkanes (e.g., methane to methanol<sup>1,2</sup>) represents one of the foremost challenges in catalysis. Despite intense focus,<sup>3–8</sup> the design of highly selective and active synthetic catalysts has been limited by overoxidation as a result of the higher reactivity of products than reactants. Conversely, it is known that enzymes readily catalyze selective partial oxidation through the formation of high-valent metal–oxo (e.g., Fe(IV)=O) species.<sup>9–12</sup> These biological catalysts have motivated the development of bioinspired molecular<sup>13–17</sup> and extended<sup>18</sup> (e.g., zeolite<sup>19</sup> or metal–organic framework<sup>20</sup>) catalysts for C–H activation.

High-valent<sup>21</sup> (e.g., Fe(IV),<sup>22,23</sup> Fe(V),<sup>24,25</sup> Mn(IV),<sup>26</sup> or Mn(V)<sup>27,28</sup>) metal–oxo species are believed to be central to selective partial oxidation, and the difficulty with which they are isolated and characterized spectroscopically<sup>23,29–32</sup> moti-

vates computational screening<sup>33,34</sup> and characterization.<sup>35–37</sup>

High-throughput computational catalyst screening can extend beyond the small number of complexes and materials that have been demonstrated experimentally to support formation of high-valent metal–oxo species, instead permitting the discovery of design rules across the periodic table.<sup>38,39</sup> Such an effort is motivated, for example, by the fact that both late (e.g., Co) and low-valent (e.g., Fe(III)–O) transition metals had been thought to be nearing the “oxo wall”,<sup>40–43</sup> but an increasing number of Co(IV)=O complexes have been recently characterized.<sup>44–46</sup> The computational study of such open-shell species with variable oxidation state is made more

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complex by the role of electron spin, an inherently quantum-mechanical property. Distinct metal–oxo spin states can have strongly differing reactivity,<sup>32,47–52</sup> with the highest rates expected for high-spin (i.e., quintet) Fe(IV)=O complexes that mimic enzymatic systems.<sup>53</sup> For each spin state and complex, stability can be as much of a challenge,<sup>16,54–57</sup> with some of the most reactive species deactivating significantly faster than less reactive species.

Large-scale computational screening<sup>33,58–70</sup> has the potential to uncover which chemical environments enable formation of stable metal–oxo species. Widely successful approaches in computational screening for heterogeneous catalysis, such as linear scaling relationships<sup>65,68,71–77</sup> or established quantum mechanical (QM) descriptors (e.g., the d-band center<sup>78–80</sup> or other frontier orbital properties<sup>61,70,81</sup>), are expected<sup>82</sup> to be limited in their capacity to describe spin-state-dependent metal–oxo formation. Indeed, even for a fixed spin state, small geometric distortions have been shown to cause large deviations in linear free energy relationships and alter the frontier orbital energies<sup>67</sup> relevant for metal–oxo formation. Noncovalent interactions relevant in single-site catalysts have also been shown to disrupt scaling relations.<sup>72,73,76,77,83–87</sup> A screen of multiple metal, spin, and oxidation states in combination with a wide range of ligands is motivated by the desire to reveal the extent to which conventional scaling rules apply or may be broken in selective partial oxidation, but the sheer combinatorial challenge of such a screen requires a different approach than standard first-principles screening.

Machine-learning (ML) property prediction models have the promise of accelerating discovery by enabling property prediction in seconds instead of the hours required by first-principles computational screening.<sup>88–92</sup> In recent years, ML has been increasingly applied to accelerate mechanism<sup>93,94</sup> or materials<sup>95–97</sup> discovery of closed-shell and/or bulk metal heterogeneous catalysts. Significant progress has also been made in the prediction of fundamentally quantum-mechanical properties of open-shell transition metal complexes, such as frontier orbital energies,<sup>98,99</sup> ionization or redox potentials,<sup>92,100,101</sup> and spin-state ordering.<sup>102–104</sup> Successful ML models have not yet been demonstrated in challenging open-shell, single-site catalysts that exhibit spin-state-dependent reactivity. It is in this area that nonlinear ML models may have the greatest promise to accelerate high-throughput screening due to the weak predictive capability of linear scaling relations, which are frequently distorted or broken in isolated, under-coordinated metal sites.<sup>67,105,106</sup>

In this work, we train ML models to predict spin-state-dependent metal–oxo formation energies in octahedral model catalysts. Using these models, we reveal unexpected structure–property trends in spin-state-dependent reactivity and the limits of the relationships between metal–oxo formation and a conventionally used QM descriptor (i.e., a frontier orbital energy). We then explore large (ca. 37 000) candidate catalyst spaces to discover wholly new complexes with unexpected combinations of oxidative stability and oxo formation energy.

## 2. CATALYSIS MODELS

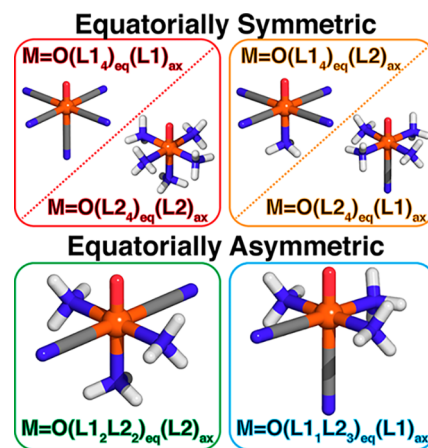
To evaluate the difficulty of machine-learning tasks for open-shell transition metal catalysis, we focus on the reaction energy for the formation of high-valent oxo species essential in C–H activation.<sup>21–28</sup> We define this oxo formation energy,  $\Delta E_{\text{oxo}}$ , as the difference in total electronic energy between a high-valent oxo species and its corresponding empty-site structure in the

same spin state with respect to a triplet oxygen reference, where the empty-site metal's oxidation state is  $n = 2$  or 3

$$\Delta E_{\text{oxo}} = E(\text{M}(n+2)=\text{O}) - E(\text{M}(n)\cdots) - \frac{1}{2}E(^3\text{O}_2) \quad (1)$$

Two data sets, the equatorially symmetric data set and the equatorially asymmetric data set, were designed to probe a wide range of effects on  $\Delta E_{\text{oxo}}$  values in octahedral, open-shell transition metal complexes. In addition to the machine-learning models, these two data sets represent 1200 new catalyst energetic evaluations carried out solely for the present work.

The equatorially symmetric (ES) data set consists of 712  $\Delta E_{\text{oxo}}$  values for complexes with one equatorial and one distal axial ligand type (Figure 1). Total complex charge varies across



**Figure 1.** Representative structures for the equatorially symmetric (top) and equatorially asymmetric (bottom) data sets, which each have up to two unique ligand types, L1 and L2 (here L1 = CN<sup>−</sup>, L2 = NH<sub>3</sub>). The metal is shown as an orange sphere, and other atoms are shown as sticks, with oxygen in red, nitrogen in blue, and carbon in gray.

catalyst models with individual ligand charge but is preserved between the oxo (M(IV)=O or M(V)=O) and the empty-site (M(II) or M(III)) complexes (M = Cr, Mn, Fe, or Co, Figure 2). Complexes are comprised of up to two distinct ligands from a pool of 29 ligands: 9 negatively charged (e.g., acac, CN<sup>−</sup>, Cl<sup>−</sup>, and pyrrole) and 20 neutral (e.g., CO, NH<sub>3</sub>, bipyridine, or furan) (Figure 2). This ligand pool was designed to broadly sample ligand fields and enable comparison to prior work.<sup>67,102,103</sup> All bidentate or tetradentate ligands can only serve as equatorial ligands.

For this data set, high-valent oxo and empty-site species were studied in their low-spin (LS), intermediate-spin (IS), and high-spin (HS) states, where defined, for both M( $n+2$ ) and M( $n$ ) oxidation states: doublet for d<sup>1</sup> Cr(V) and d<sup>3</sup> Cr(III), singlet/triplet for d<sup>2</sup> Cr(IV)/Mn(V) and d<sup>4</sup> Cr(II)/Mn(III), doublet/quartet for d<sup>3</sup> Mn(IV)/Fe(V) and d<sup>5</sup> Mn(II)/Fe(III), singlet/triplet/quintet for d<sup>4</sup> Fe(IV)/Co(V) and d<sup>6</sup> Fe(II)/Co(III), and doublet/quartet for d<sup>5</sup> Co(IV) and d<sup>7</sup> Co(II). The final data set is broadly balanced across each of these metals, oxidation states, and spin states (Supporting Information Figure S1).

The equatorially asymmetric (EA) data set consists of 488  $\Delta E_{\text{oxo}}$  values constructed from a smaller pool of 18 ligand types, 7 of which overlap with the ES data set (Figures 1 and 2





of five and a maximum tree depth of two, which were selected to reveal dominant trends.

## 4. RESULTS AND DISCUSSION

**4.1. Data Sets and Model Performance.** The range of  $\Delta E_{\text{oxo}}$  values observed in the 712 equatorially symmetric and 488 equatorially asymmetric data set complexes is comparable, from as low as  $-105$  kcal/mol to as high as  $75$  kcal/mol (Supporting Information Figure S9). Across both data sets Cr and Mn complexes in all spin states have the most exothermic  $\Delta E_{\text{oxo}}$  values, midrow Fe or Co complexes are more variable, and  $\Delta E_{\text{oxo}}$  is exclusively endothermic for the later Ni and Cu transition metals. Although the metal is a strong determinant in  $\Delta E_{\text{oxo}}$  favorability, the large variation within each metal motivates the training of ML models and further evaluation of chemical trends.

We start by extending our prior approach of employing RAC features with KRR models in the prediction of octahedral spin-splitting energetics,<sup>101</sup> redox or ionization potential,<sup>92,101</sup> metal–ligand bond length,<sup>101</sup> and frontier orbital energetics.<sup>98</sup> Assessing the degree to which this RAC/KRR approach is also predictive for  $\Delta E_{\text{oxo}}$  will (i) identify if additional complexities arise for the learning task of spin-state-dependent catalyst energetics and (ii) determine if catalytic structure–property relationships differ from those obtained previously for other properties. Thus, we repeat our approach and make comparisons throughout to both model performance and observations from prior work.<sup>92,98,101</sup> The range of reaction energies in our ML model training data is comparable to those we studied<sup>103</sup> in ML models for spin-splitting energy (from  $-55$  to  $90$  kcal/mol, Supporting Information Figure S10).

Feature selection from the full RAC feature set (ca. 150 descriptors) improves KRR model prediction errors<sup>98,101</sup> and provides insight into the most important descriptors for property prediction. Indeed, train/test mean absolute errors (MAEs) with the full set of RACs ( $7.5$  and  $9.5$  kcal/mol) for the ES data set are reduced after RF-RFA feature selection (see section 3). A model trained on only the 22 features selected with RF-RFA has significantly lower train and test MAEs ( $2.2$  and  $5.5$  kcal/mol), consistent with prior work<sup>98</sup> (Figure 3 and Supporting Information Figure S11 and Table S7). The full RAC/KRR performance is slightly better for the smaller EA

data set (MAEs, train  $2.3$  and test  $6.5$  kcal/mol), but it is similarly improved after feature selection to 14 features (MAEs, train  $1.5$  and test  $4.3$  kcal/mol, Figure 3 and Supporting Information Figure S11, Table S8, and Text S3).

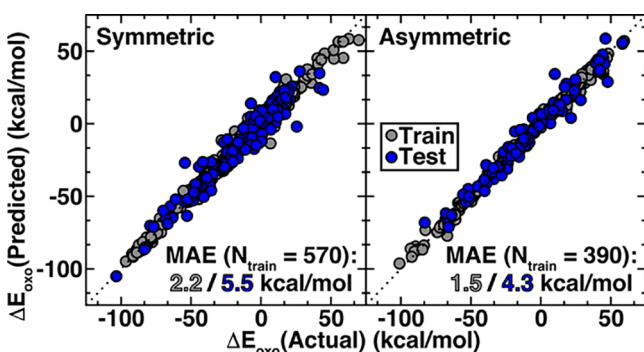
Decomposing errors by metal, oxidation state, and ligand identity provides insight into whether the RF-RFA/KRR model generalizes more poorly to specific metal–oxo complexes (Supporting Information Figures S12 and S13). The ES errors are lower for  $M(\text{IV})=\text{O}$  complexes and higher for  $M(\text{V})=\text{O}$  complexes absent from the EA data set (Supporting Information Figures S12 and S13). Nearly all of the high ( $>10$  kcal/mol, ca. 30 cases) train or test set errors in the ES data set are indeed  $M(\text{V})=\text{O}$  complexes with strong-field distal axial ligands (e.g.,  $\text{NMe}_3$ ,  $\text{CO}$ ,  $\text{pisc}$ ) or negatively charged (e.g.,  $\text{Cl}^-$ ,  $\text{CN}^-$ ,  $\text{acac}$ ,  $\text{pyrrole}$ ) equatorial ligands. Across both data sets, test errors are generally lower for Mn and Fe in comparison to earlier or later metals (Supporting Information Figures S12 and S13). Comparison of errors across the EA data set ligand arrangements indicates slightly higher errors for the more asymmetric  $M=\text{O}(\text{L}_1\text{L}_2)_\text{eq}(\text{L}_1)_\text{ax}$  configuration, likely due to its lower abundance in the data set (37%, Supporting Information Figures S13 and S14).

Over both the ES and EA data sets, test MAEs of  $4.3$  and  $5.8$  kcal/mol are somewhat larger than the smallest we observed for spin-splitting energies<sup>101,103</sup> (ca.  $1$ – $3$  kcal/mol) but roughly comparable to the  $4$  kcal/mol MAEs we observed for gas-phase ionization potential,<sup>101</sup> redox potential predictions,<sup>101</sup> or frontier orbital energies.<sup>98</sup> Given the wide diversity of ligand charges, structures, and metals as well as the fact that up to three  $\Delta E_{\text{oxo}}$  values are predicted for each complex, the performance of the present KRR models is unexpectedly good. Spin-state-dependent reaction energetics of metal–oxo formation do not appear to be more challenging to predict than other properties of transition metal complexes.

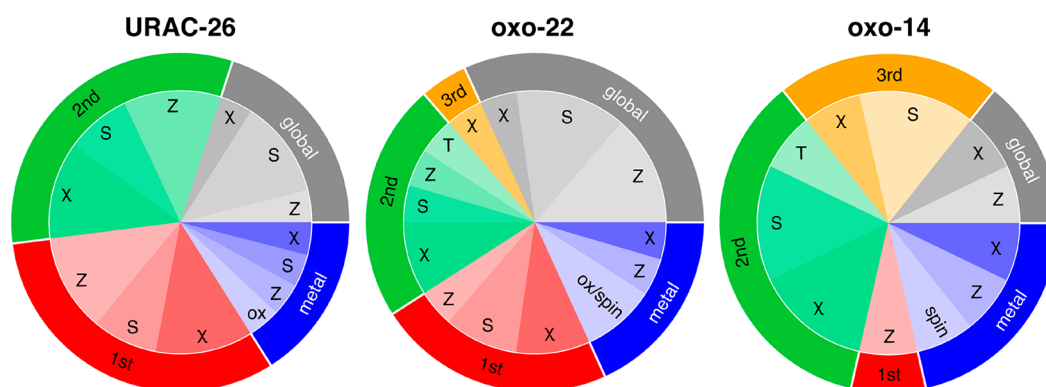
### 4.2. Features that Influence Oxo Formation Energies.

Beyond test set performance, comparison of RFA-selected features (oxo-22 for ES, oxo-14 for EA) reveals the length scale and character of molecular features that drive  $\Delta E_{\text{oxo}}$  values (Figure 4, Supporting Information Tables S7 and S8). As in previous work,<sup>101</sup> the five heuristic atomic properties we correlate in approximately 150 RACs are (i)  $I$ , the identity, (ii)  $T$ , topology (i.e., connectivity), (iii)  $\chi$ , Pauling electronegativity, (iv)  $S$ , the covalent radius, and (v)  $Z$ , the nuclear charge. These quantities are derived from products and differences of atomic properties on the molecular graph that we obtain either over the whole molecule (i.e., global features) or by counting bond paths outward from the metal center (e.g., first, second, or third coordination-sphere RACs, see Supporting Information Text S2). We compare the oxo-selected sets to a 26 RAC feature set (URAC-26) that was selected on spin-splitting data but has been shown to have the best balanced performance on ionization potential, redox potential, and bond length property prediction<sup>92,101</sup> (Figure 4). As could be expected from our preliminary analysis of the  $\Delta E_{\text{oxo}}$  data set, properties of the metal, oxidation state, and spin are essential (ca. 20–25% of all features) and of comparable weight to those in URAC-26 (Figure 4, Supporting Information Table S9).

We previously found metal-coordinating atoms in the first coordination sphere to be important in URAC-26 and related metal-local feature sets for prediction of spin splitting,<sup>103</sup> an observation that holds for the ES oxo-22 feature set but not for the EA oxo-14 feature set (Figure 4). Differences in feature



**Figure 3.** KRR model  $\Delta E_{\text{oxo}}$  predictions compared to the DFT-calculated values for the equatorially symmetric (left) and equatorially asymmetric (right) data sets with RACs obtained from feature selection on each set, respectively. Number of training points used after the 80/20 train/test split is indicated in the inset along with the train (gray circles) and test (blue circles) mean absolute error (MAE) in kcal/mol. A black dotted parity line is also shown.



**Figure 4.** Pie charts of the URAC (26 features, left) features selected in prior work<sup>92,101</sup> compared to features selected by RF-RFA for oxo formation on the equatorially symmetric data set (22 features, middle) or the equatorially asymmetric data set (14 features, right). Features are grouped by the most distant atoms present: metal in blue, first coordination sphere in red, second coordination sphere in green, third coordination sphere in orange, or global features in gray. Within each distance category, the property (i.e.,  $\chi$ , S, T, or Z) is also indicated, and oxidation state (ox) and spin are assigned as metal-local properties.

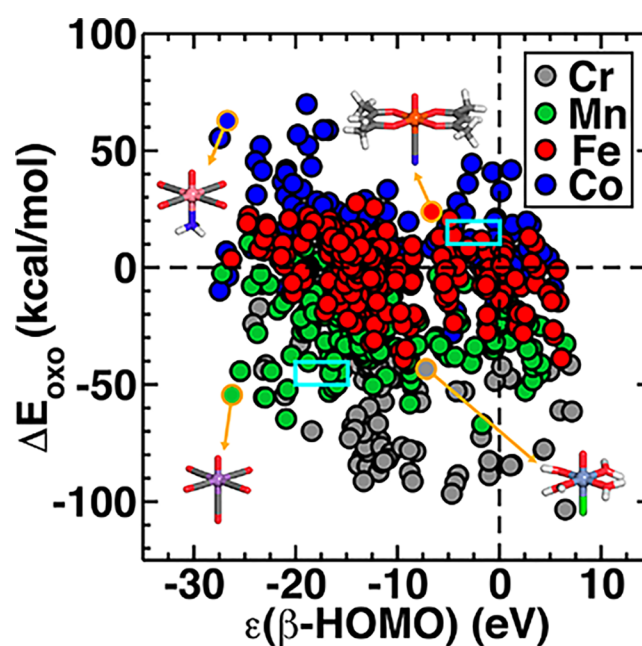
selection over the ES and EA data sets can be attributed to the reduced size and variation of chemical composition for the EA data set (Supporting Information Text S4, Figures S15–S18, and Tables S10 and S11). Overall, feature analysis is broadly useful for interpreting data sets and ML models, but transferability of feature sets across different KRR prediction models is generally observed (Supporting Information Tables S9 and S12 and Figure S19).

The  $\Delta E_{\text{oxo}}$ -selected features distinguish themselves from URAC-26 in their increased weight (ca. 35% vs 20%) of third coordination sphere or global features (Figure 4). Across the ES and EA data sets, these differences appear to be most critical to encoding through-bond ligand variations, as none of these metal-distant atoms are sufficiently proximal to form noncovalent interactions with the  $\text{M(IV/V)=O}$  species.<sup>67,85–87</sup> This feature analysis reveals oxo formation design rules:  $\Delta E_{\text{oxo}}$  is affected not just by metal identity but also relatively metal-distant, through-bond electronic<sup>101</sup> (i.e.,  $\chi$  or Z) ligand functionalization, expanding upon previous experimental observations.<sup>128</sup>

#### 4.3. Relation to QM Descriptors for Oxo Formation.

To accelerate and simplify computational screening, the  $\beta$ -HOMO level of  $\text{Fe(II)}$  complexes has been used as an estimate of the favorability of quintet  $\text{Fe(IV)=O}$  complex formation as long as this level has significant d character.<sup>70</sup> If this approximation holds<sup>70</sup> it provides an intuitive rationale that the metal must be easily oxidized for the high-valent species to form. Across the ES data set, most  $\beta$ -HOMO levels of  $\text{Fe(II)}$  complexes indeed have significant d character, with a few exceptions for conjugated ligands (e.g., pisc, bifuran, and cyanopyridine), but this observation holds less well across metals (Supporting Information Figures S20 and S21 and Table S13). A large range of  $\beta$ -HOMO level values is observed from around  $-30$  to  $+8$  eV, where the unphysically positive  $\beta$ -HOMO levels correspond to cases where high negative complex charge leads to poorly bound electrons at the hybrid DFT level of theory (Supporting Information Figure S22).<sup>129,130</sup> Over the full ES data set,  $\Delta E_{\text{oxo}}$  and the  $\beta$ -HOMO level correlate only weakly overall and within a single metal, regardless of d character in the orbital (Figure 5).

As an example of the ease with which this  $\beta$ -HOMO/ $\Delta E_{\text{oxo}}$  design rule may be broken,  $S = 0$   $\text{Co(V)=O}(\text{CO})_4(\text{NH}_3)$  and  $\text{Mn(V)=O}(\text{CO})_5$  have comparably deep  $\beta$ -HOMO levels (from ca.  $-26$  to  $-27$  eV), but  $\Delta E_{\text{oxo}}$  is unfavorable by the



**Figure 5.** Empty-site structure  $\beta$ -HOMO level (in eV) vs  $\Delta E_{\text{oxo}}$  (in kcal/mol) for the 712 structures in the equatorially symmetric set, colored by metal identity. Four representative structures are shown as insets: (top left)  $S = 0$   $\text{Co(V)=O}(\text{CO})_4(\text{NH}_3)$ , (top right)  $S = 1/2$   $\text{Fe(V)=O}(\text{acac})_2(\text{CN}^-)$ , (bottom left)  $S = 0$   $\text{Mn(V)=O}(\text{CO})_5$ , (bottom right)  $S = 1$   $\text{Cr(IV)=O}(\text{H}_2\text{O})_4(\text{Cl}^-)$ . Atoms are colored as follows: purple for Mn, pink for Co, light blue for Cr, red for O, blue for N, white for H, gray for C, and green for Cl. Cyan rectangles for Zone 1 (bottom, left) and Zone 2 (top, right) are also shown.

same magnitude for  $\text{Co(V)=O}$  (55.4 kcal/mol) as it is favorable for  $\text{Mn(V)=O}$  ( $-54.4$  kcal/mol, Figure 5). Weaker field ligands (e.g.,  $S = 1/2$   $\text{Fe(V)=O}(\text{acac})_2(\text{CN}^-)$  and  $S = 1$   $\text{Cr(IV)=O}(\text{H}_2\text{O})_4(\text{Cl}^-)$ ) correspond to shallower  $\beta$ -HOMO levels (e.g., from  $-7$  to  $-9$  eV), but  $\Delta E_{\text{oxo}}$  values span a large range of endothermic and exothermic values for these complexes ( $\text{Fe(V)=O}$  23.9 kcal/mol,  $\text{Cr(IV)=O}$  46.5 kcal/mol, Figure 5). Even within a fixed metal, variations of tens of electronvolts of the  $\beta$ -HOMO level can be observed at fixed  $\Delta E_{\text{oxo}}$  values or 50–100 kcal/mol variations of  $\Delta E_{\text{oxo}}$  at fixed  $\beta$ -HOMO level; these observations suggest the possibility of orthogonally tuning frontier orbital energies (e.g., for resting

state complex oxidative stability) and oxo formation energies (e.g., for reactivity).

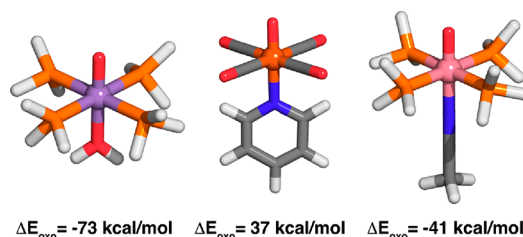
Given the weak correlation between the  $\beta$ -HOMO level and  $\Delta E_{\text{oxo}}$ , we developed a separate ML model to predict the  $\beta$ -HOMO level. We trained RF-RFA KRR models on the ES data set to obtain a test MAE of 0.44 eV that was somewhat higher than our prior frontier orbital energy models<sup>98</sup> (Supporting Information Figures S23–S26). Analyzing the 33 selected features for the RF-RFA  $\beta$ -HOMO level KRR model reveals the physical basis for the weak correlation between the properties (Figure 4 and Supporting Information Figure S27 and Table S14). The 33 features selected for  $\beta$ -HOMO level prediction are predominantly global (76%) and third coordination sphere (21%) properties that now include an enhanced dependence on the coordination of atoms in the structure (i.e., *T* or *I* RACs), with only weak contributions from the metal or second coordination sphere ( $\chi$  only) and none from the first coordination sphere. Given the strong role of complex size and ligand character in determining the  $\beta$ -HOMO level, ligand chemistry or metal identity would need to be fixed to establish a correlation to oxo formation energies.

#### 4.4. Enumeration of a Theoretical Catalyst Space.

From the 29 ligands originally considered in the ES data set, there is a theoretical space of 9860 transition metal complexes formed from all allowed combinations of metals, oxidation and spin states, and ligands. Although this space is inherently interpolative in nature, only 7.2% (712 points: 570 in train, 142 in test) of this compound space was used during model construction. Failed calculations account for 9.5% (938 points) of the space, but the remaining 83.3% (8210 points) are complexes for which no DFT calculation had been attempted. We switch from KRR to ANN models for the full space enumeration due to our experience<sup>103</sup> that ANNs generalize better than KRR models at the cost of being harder to interpret (Supporting Information Figures S28–S34).

The  $\Delta E_{\text{oxo}}$  and  $\beta$ -HOMO level value ranges across the ML-model-enumerated theoretical space suggest wide coverage of the full range of  $\beta$ -HOMO level values within a single metal and significant overlap across metals for  $\Delta E_{\text{oxo}}$  values (Supporting Information Figure S35). To confirm these observations, we obtained DFT validation results on a set of 277 previously unseen complexes obtained from ML model minimum, median, or maximum  $\Delta E_{\text{oxo}}$  values for each metal (Supporting Information Figures S31, S32, and S35 and Table S15). These new DFT results further reinforce the observation that  $\Delta E_{\text{oxo}}$  values and the  $\beta$ -HOMO level can be tuned independently. As an example of this design exception, a 14 eV variation in  $\beta$ -HOMO level is observed with limited change in  $\Delta E_{\text{oxo}}$  for two Co(V)=O complexes: Co(V)=O(acac)<sub>2</sub>(Cl<sup>−</sup>) ( $\beta$ -HOMO = −6.2 eV,  $\Delta E_{\text{oxo}}$  = 47.3 kcal/mol) vs Co(V)=O(NH<sub>3</sub>)<sub>4</sub>(OMe<sub>2</sub>) ( $\beta$ -HOMO = −20.6 eV,  $\Delta E_{\text{oxo}}$  = 43.0 kcal/mol) (Figure 5 and Supporting Information Figure S36).

Because the points selected for validation were  $\Delta E_{\text{oxo}}$  extrema in the ML-model-interpolated space, many such points exceed the bounds of the original 712 DFT points in the ES data set and are extrapolative in property space. For example, singlet Mn(V)=O complexes with equatorial phosphine ligands and weak-field (e.g., H<sub>2</sub>O −73 kcal/mol or misc −68 kcal/mol) axial ligands have even more favorable  $\Delta E_{\text{oxo}}$  values than had been observed (singlet Mn(V)=O(pyrrole)<sub>4</sub>(H<sub>2</sub>O) −67 kcal/mol) before (Figure 6). Although most Fe(IV)=O complexes have favorable  $\Delta E_{\text{oxo}}$  values, three new singlet Fe(IV)=O(CO)<sub>4</sub>(ax) complexes (e.g., ax =



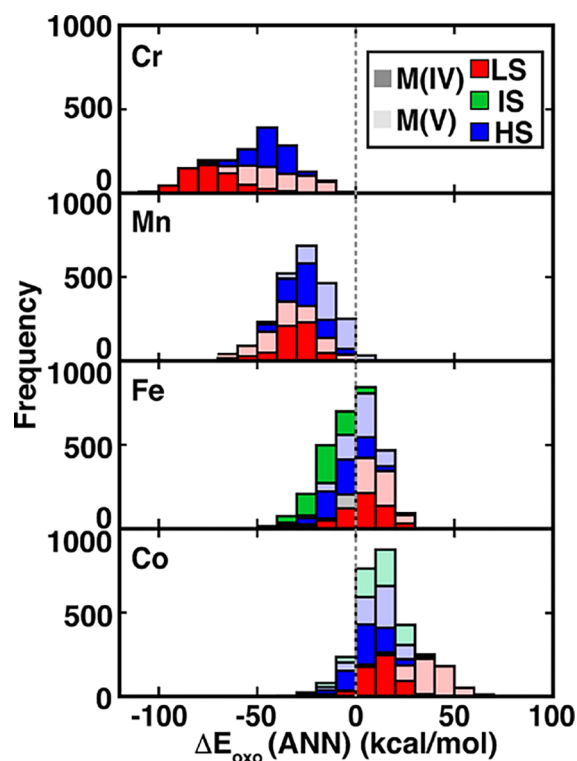
**Figure 6.** Three extreme points from the new data set with their  $\Delta E_{\text{oxo}}$  values: singlet Mn(V)=O(PH<sub>3</sub>)<sub>4</sub>(H<sub>2</sub>O) (left), singlet Fe(IV)=O(CO)<sub>4</sub>(pyridine) (middle), and singlet Co(V)=O(PH<sub>3</sub>)<sub>4</sub>(CH<sub>3</sub>CN) (right). Atoms in the stick structures of each lead compound are as follows: purple for Mn, dark orange for Fe, pink for Co, blue for N, gray for C, red for O, white for H, and orange for P.

pyridine 37 kcal/mol) exceed the original ES data set's highest  $\Delta E_{\text{oxo}}$  value compound (Fe(IV)=O(misc)<sub>5</sub> 28 kcal/mol), reinforcing equatorial CO as unfavorable for oxo formation across a range of axial ligands (Figure 6). The quintet Co(V)=O(CN<sup>−</sup>)<sub>4</sub>(H<sub>2</sub>O) with the most favorable  $\Delta E_{\text{oxo}}$  value (−28 kcal/mol) in the ES data set has also been exceeded by as much as 13 kcal/mol in the likely ground state singlet Co(V)=O(PH<sub>3</sub>)<sub>4</sub>(CH<sub>3</sub>CN) (−41 kcal/mol, Figure 6). Analysis of simultaneously deep  $\beta$ -HOMO levels (<−15 eV) and favorable  $\Delta E_{\text{oxo}}$  values (<−50 kcal/mol) in the new DFT data set reveals that in addition to singlet Mn(V)=O complexes with N-coordinating ligands that we observed in the original ES data set, we now confirm that equatorial PH<sub>3</sub> and weak-field axial ligands (i.e., singlet Mn(V)=O(PH<sub>3</sub>)<sub>4</sub>(ax) where ax = misc, Cl<sup>−</sup>, furan, or water) populate this zone (Supporting Information Table S16).

Across the ANN-enumerated space, the ranges of metal-dependent  $\Delta E_{\text{oxo}}$  values agree with the observation<sup>131</sup> that increasing d-electron count makes  $\Delta E_{\text{oxo}}$  increasingly unfavorable as M–O  $\pi^*$  antibonding orbitals<sup>132,133</sup> become occupied (e.g., Fe or Co in Figure 7). The oxo formation energies for Cr and Mn complexes remain exothermic with an exceedingly small fraction (2.8%, 33 of 1160) of strong-field (e.g., CO) Mn(V) endothermic complexes.

Although spin-state dependence is expected<sup>85</sup> in Fe(IV)=O and its isoelectronic analogue, Co(V)=O, with more favorable  $\Delta E_{\text{oxo}}$  values for HS and IS than LS, the enumerated data set reveals deeper chemical trends (Figure 7). The Fe(IV)=O complexes have the highest spin-state dependence when CO or CN<sup>−</sup> are equatorial ligands, as is expected (favorable HS, IS; unfavorable LS, Supporting Information Figure S37). Weakening the equatorial or axial ligand (e.g., to H<sub>2</sub>O or Cl<sup>−</sup>) lessens this spin-state-dependent effect (Supporting Information Figures S37 and S38). Little spin-state dependence is evident in the Fe(V)=O complexes, which generally have similar LS and HS  $\Delta E_{\text{oxo}}$  values (Supporting Information Figure S39). Starting from a weak equatorial ligand (e.g., water) in Fe(IV)=O without any apparent spin-state dependence and introducing strong-field ligands recovers moderate spin-state dependence on  $\Delta E_{\text{oxo}}$  with favorable IS but unfavorable LS or HS values (Supporting Information Figures S37 and S38). Although isoelectronic Co(V)=O shows variation in the magnitudes of LS, IS, and HS  $\Delta E_{\text{oxo}}$  values, nearly all are unfavorable (Supporting Information Figure S40). Interestingly, the combination of strong-field equatorial CO ligands with a weak-field (e.g., water) axial ligand that was observed to lessen Fe(IV)=O spin-state



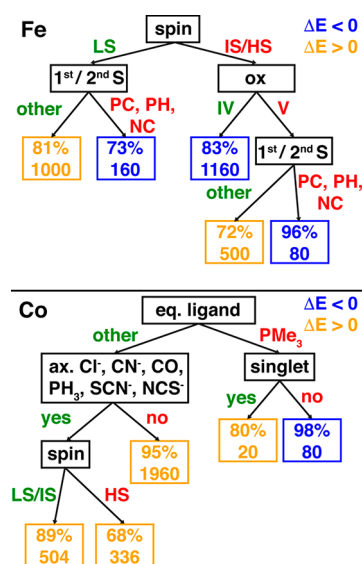


**Figure 7.** Distribution of oxo formation energies (in kcal/mol, bin size 10 kcal/mol) as predicted by the ANN for the 9860 equatorially symmetric complex space. Unnormalized counts are shown on the y axis, and histogram is colored by spin (red for low spin, LS, green for intermediate spin, IS, and blue for high spin, HS). The stacked histogram is shaded by oxidation state, with oxidation state +5 complexes represented by translucent coloring and oxidation state +4 complexes represented by opaque coloring.

dependence is instead predicted to strengthen Co(V)=O complex spin-state dependence (Supporting Information Figures S37 and S40).

To broadly uncover the governing factors in  $\Delta E_{\text{oxo}}$  values, we carried out decision tree analysis on all enumerated Fe and Co ES-compatible complexes. Indeed, spin state is the largest driver for favorability, with most IS or HS complexes having favorable  $\Delta E_{\text{oxo}}$  values (Figure 8). For the less stable high-spin Fe(V) complexes, strong-field ligand atoms in the equatorial plane first and second coordination spheres (e.g., P–C, P–H, and N–C first/second S) result in favorable  $\Delta E_{\text{oxo}}$  values in the majority of cases (96%, Figure 8). Although such strong-field ligands are likely to bias toward LS ground states, this composition of the metal environment also is correlated with favorable  $\Delta E_{\text{oxo}}$  values for both Fe(IV)=O and Fe(V)=O low-spin complexes (73%, Figure 8).

For Co, trends are less clear, likely because so few Co(IV/V)=O complexes are stable (Figures 7 and 8). From the decision tree analysis, IS and HS  $\text{PMe}_3$  Co–oxo complexes are likely to have favorable  $\Delta E_{\text{oxo}}$  values, but these are unlikely to be ground states (Figure 8). Indeed, from all interpolated complexes, only 11.8% (342 of 2900) are predicted by the ANN to have favorable  $\Delta E_{\text{oxo}}$  values, and most have high- or intermediate-spin Co complexed with equatorial pnictogen atoms (108 N and 127 P) and axial strong-field coordinating atoms (129 C, 98 N, and 43 P). Although ligand field arguments suggest that HS or IS Co complexes would be rare, observations from our enumeration are consistent with the



**Figure 8.** Decision tree analysis for essential descriptors of favorability for Fe (top) and Co (bottom) ANN-predicted  $\Delta E_{\text{oxo}}$  binary decision tree divided by favorable ( $\Delta E < 0$ , blue) and unfavorable ( $\Delta E > 0$ , orange) for up to three levels total. Each leaf shows the percent of data corresponding to the case and total number of interpolated-space ANN values that correspond to each leaf.

recent isolation and spectroscopic characterization of a metastable, N-coordinating complex, quartet [(13-TMC)Co(IV)=O]<sup>2+</sup> (13-TMC is 1,4,7,10-tetramethyl-1,4,7,10-tetraazacyclotridecane).<sup>134</sup>

**4.5. ML-Driven Catalyst Space Exploration.** Inspired by the possibility of orthogonal empty-site  $\beta$ -HOMO level and  $\Delta E_{\text{oxo}}$  tuning, we set out to search a larger design space of transition metal complexes. Our aim was to populate zones of empty-site  $\beta$ -HOMO level and  $\Delta E_{\text{oxo}}$  that were underpopulated in our original screen and that generally defy chemical intuition. For example, equatorially symmetric model catalysts could defy expectations by having (i) shallow empty-site  $\beta$ -HOMO levels, i.e., low ionization potentials, in combination with unfavorable  $\Delta E_{\text{oxo}}$ , or (ii) deep empty-site  $\beta$ -HOMO levels, i.e., high ionization potentials, and favorable  $\Delta E_{\text{oxo}}$ . In either case, we expect ANN-accelerated screening to reveal what catalyst model chemistries are most suited to breaking the relationship between resting state ionization potential and oxo formation favorability.

To carry out exploration beyond interpolative enumeration, we expanded the possible design space to a total of 56 different ligands (39 monodentate, 17 multidentate) in ES-compatible combinations, i.e., 27 more than the 29 originally included in the ES data set (Supporting Information Figure S41 and Table S17). Although several ligands were previously in the EA data set, we incorporated common ligands (e.g., tetraphenylporphyrin, phthalocyanine, bipyrimidine) and functionalizations of ES ligands (e.g., phendione, phenacac, mebpy). These new ligands enlarge the design space to 37 128 complexes (Supporting Information Table S18).

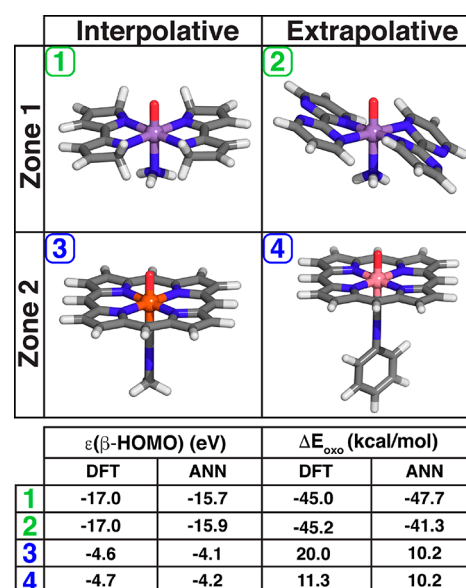
To accelerate discovery we combined our independent ANNs with a genetic algorithm (GA) optimization that we previously demonstrated for designing transition metal complexes with near-degenerate spin-splitting energies<sup>102</sup> and targeted band gaps.<sup>98</sup> For the present GA to optimize  $\beta$ -HOMO levels and  $\Delta E_{\text{oxo}}$  simultaneously, we employed a composite fitness function with distance awareness to avoid

points where the ANN will lack predictive power (Supporting Information Text S1, Figure S42, and Table S19). We then targeted two distinct regions of property space. For the first targeted zone (Zone 1), we selected  $\Delta E_{\text{oxo}}$  between  $-40$  and  $-50$  kcal/mol along with a simultaneous empty-site  $\beta$ -HOMO level between  $-20$  and  $-15$  eV. Five ES complexes (train 5, test 0) are within this range, all of which are singlet  $\text{Mn(V)=O}$  coordinated in the equatorial plane by nitrogen atoms (e.g., ammonia or bpy see Figure 5). For the second targeted zone (Zone 2), we selected  $\Delta E_{\text{oxo}}$  between  $10$  and  $20$  kcal/mol along with a simultaneous empty-site  $\beta$ -HOMO level between  $-5$  and  $0$  eV. Six ES data points (train 5, test 1), two  $\text{Fe(IV)=O}$  and four isoelectronic  $\text{Co(V)=O}$  complexes, with negatively charged equatorial ligands reside in this zone. A GA approach is expected to help enrich both zones, where Zone 1 corresponds to a likely desired, stable catalyst, whereas Zone 2 corresponds to catalysts with less favorable energetics than would be predicted by the  $\beta$ -HOMO descriptor.

The composite-objective distance-controlled GA was run 10 times each for Zones 1 and 2 (Supporting Information Figures S43 and S44). For each zone, 50 leads (out of 137 total leads) generated from the runs were selected for further study with DFT. The majority of such structures correspond to unstable molecules at the DFT level, explaining why they were sparsely populated in our original DFT data set: only approximately 25% of the data (Zone 1: 13 leads, Zone 2: 15 leads) passed our standard electronic or geometric structure<sup>135</sup> checks (Supporting Information Table S20).

For Zone 1, i.e., favorable  $\Delta E_{\text{oxo}}$  and high ionization potential, four new DFT lead compounds fully satisfy both criteria, nearly doubling the data we had in this range of properties, and the remaining complexes all satisfy the  $\beta$ -HOMO level range but fall narrowly below or above the target  $\Delta E_{\text{oxo}}$  (Supporting Information Table S20). All four leads that fully satisfy the Zone 1 criteria are again singlet  $\text{Mn(V)=O}$  model catalysts, and three have the equatorial 2,2'-bipyrimidine ligand (bpym) that we introduced in the extended GA design space with a range of intermediate-field axial ligands (e.g., with a new imidazoline axial ligand, Figure 9 and Supporting Information Figure S41 and Table S17). Thus, DFT-validated lead compounds from the GA maintain and strengthen the observation that N-coordinated singlet  $\text{Mn(V)=O}$  model catalysts will have favorable  $\Delta E_{\text{oxo}}$  and deep  $\beta$ -HOMO levels. Although the remaining 9 leads fall outside the Zone 1 target for  $\Delta E_{\text{oxo}}$ , five cases are doublet  $\text{Cr(V)=O}$  complexes that still satisfy the deep  $\beta$ -HOMO level target (Supporting Information Table S20). The doublet  $\text{Cr(V)=O}(\text{CN-pyr})_4(\text{misc})$  is nearest to Zone 1 (DFT  $\Delta E_{\text{oxo}} = -32$  kcal/mol,  $\beta$ -HOMO =  $-16.5$  eV), but the ANN predictions are eroded here due to the limited number of similar compounds in the training data. Overall, these DFT leads enhance data density in and around Zone 1. These complexes could be the focus of future computational study in selective partial oxidation (e.g., to understand the potential effect of deep  $\beta$ -HOMO levels).

For Zone 2, i.e., the unexpected combination of unfavorable  $\Delta E_{\text{oxo}}$  and shallow  $\beta$ -HOMO level, two leads fall fully within both ranges after DFT validation and an additional four satisfy one of the two criteria. The two Zone 2 leads are quintet  $\text{Fe(IV)=O}$  or quartet  $\text{Co(IV)=O}$  porphyrin complexes with axial functionalized-isocyanide ligands (Figure 9). No prior training data in this zone had been  $\text{Co(IV)=O}$ , emphasizing the ability of this approach to discover new chemistry. All four leads that satisfy one of the two Zone 2 ranges are also



**Figure 9.** Leads from the GA exploration, as predicted by the ANN and validated by DFT, that are observed for the two targeted zones 1 and 2 as described in the main text. Leads using ligands from the original equatorially symmetric ligand pool are labeled as “interpolative”, and leads using ligands from a distinct, expanded ligand pool are labeled as “extrapolative”. Atoms in the stick structures of each lead compound are as follows: purple for Mn, orange for Fe, pink for Co, blue for N, gray for C, red for O, and white for H.

$\text{Co(IV)=O}$  complexes, e.g., (i) quartet complexes with equatorial acac ligands that have slightly more favorable DFT  $\Delta E_{\text{oxo}}$  values ( $1\text{--}4$  kcal/mol) than predicted by the ANN and (ii) a quartet complex with neutral furan equatorial ligands in combination with an expanded design space phosphorine axial ligand that has a weakly positive  $\beta$ -HOMO ( $1$  eV, Supporting Information Table S20). For the remaining cases that failed to meet both Zone 2 criteria, most have weakly more favorable  $\Delta E_{\text{oxo}}$  values combined with less favorable (i.e., positive)  $\beta$ -HOMO levels. These complexes represent cases where design based on  $\beta$ -HOMO level would fail and could therefore be used in future study to understand the limitations of QM descriptors for reactions involving oxo intermediates.

## 5. CONCLUSIONS

We trained ML models capable of predicting oxo formation reaction energies across a range of first-row metals, oxidation states, and spin states. With feature-selected ML models, we achieved set-aside test mean absolute errors of  $4\text{--}5$  kcal/mol across a range of ligand orientations. These comparable errors to other properties of open-shell transition metal complexes (e.g., ionization potential) suggest that spin-state-dependent catalyst structure–property relationships are no more challenging a learning task. Using feature selection, we observed that the most important features for predicting oxo formation energies were more nonlocal in nature than for spin-state ordering, likely due to enhanced importance of through-bond electronic effects in determining the stability of metal–oxo intermediates. We then used an ML model to enumerate the space spanned by our ligand set, enabling widespread determination of spin- and metal-dependent trends. The enumeration also revealed unexpected metal/ligand-property relationships, such as reduced spin-state dependence in  $\text{Fe(IV)=O}$  complexes with strong-field equatorial ligands



and weak-field axial ligands that instead increased spin-state dependence for Co(V)=O complexes.

In contrast with earlier computational screens where the  $\beta$ -HOMO level of an M(II) complex could be shown to predict its M(IV)=O stability, we observed only weak correlation over our DFT data sets. Full ANN enumeration of both properties in this space revealed opportunities to break this design rule by producing oxidatively stable complexes that favorably form oxo intermediates. We then expanded the search space to over 37 000 catalysts by introducing new ligands and used a multicomponent GA to discover candidate catalysts that defy expectations but could still be confidently predicted by our ANN models. Using this approach, we identified both favorable oxo formation energies for oxidatively stable complexes (i.e., a typical target for a catalyst screen) and unfavorable oxo formation energies for oxidatively unstable complexes (i.e., a combination unexpected by intuition). This approach doubled the number of DFT hits that satisfied these constraints, including complexes distinct from what had been seen before in DFT training data. These observations point to the opportunities for ML-model driven discovery to both identify desirable catalysts where Edisonian approaches have failed and to find ways to break rules when known exceptions are limited. Next steps beyond the current approach will be to consider multiple reaction steps, transition states, and through-space interactions neglected thus far in our representations.

## ■ ASSOCIATED CONTENT

### ■ Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscatal.9b02165.

Ligands from both data sets; statistics of data sets separated by metal, oxidation state, and spin state; criteria used for geometry checks, failure modes, and counts for the equatorially symmetric set; hyperparameters and training information for KRR and ANN models; random forest preordered recursive feature addition (RF-RFA) for both data sets; distribution of oxo formation energies and comparisons to previous work on spin-splitting energies; nonfeature-selected KRR model performance; features selected on both data sets and comparisons between feature sets; comparison of M06-L/def2-TZVP vs B3LYP/LACVP\* oxo formation energies sorted by connecting atom identity; structures of endothermically biased oxo formation energies by M06-L/def2-TZVP; train and test performance of KRR models for M06-L/def2-TZVP data; KRR errors decomposed by metal, oxidation state, and/or arrangement; d-orbital character of empty-site  $\beta$ -HOMO in quintet Fe complexes and full equatorially symmetric set; empty-site  $\beta$ -HOMO level vs oxo structure energetic HOMO level; comparison of previously published frontier orbital energetics data set to equatorially symmetric data set; application of previously published model for frontier orbital energetics on equatorially symmetric data; full enumeration predictions by the feature-selected KRR/ANN models; histogram of deviations between the feature-selected KRR and ANN models; ANN enumerated oxo formation energy vs  $\beta$ -HOMO level; placement of oxo formation energy and  $\beta$ -HOMO level for a 277 molecule validation set; plot of absolute errors for  $\beta$ -HOMO level

and oxo formation energy; expanded ligand space used in the genetic algorithm (GA) exploration; latent distance cutoff calibration; evolution of the GA by generation for both zones; average distance and diversity over the 10 replicates for each targeted zone; examples for feature space vs latent space distances (PDF)

Structures from the equatorially symmetric data set; 277 molecule DFT validation data set; GA lead compounds from Zones 1 and 2; raw electronic energies for the equatorially symmetric data set (B3LYP/LACVP\* and M06L/def2-TZVP), equatorially asymmetric data set, 277 molecule DFT validation data set, and GA lead compounds from Zones 1 and 2; RAC featurization for equatorially symmetric and equatorially asymmetric data sets (ZIP)

## ■ AUTHOR INFORMATION

### Corresponding Authors

\*E-mail: rgetman@g.clemson.edu. Phone: 864-656-5423.

\*E-mail: hjkulik@mit.edu. Phone: 617-253-4584.

### ORCID

Aditya Nandy: 0000-0001-7137-5449

Jon Paul Janet: 0000-0001-7825-4797

Chenru Duan: 0000-0003-2592-4237

Heather J. Kulik: 0000-0001-9342-0191

### Notes

The authors declare no competing financial interest.

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